1

WHAT IS CLAIMED IS:

2 transduced in a batch.

1	1. A method of identifying a gene or genes associated with a selected				
2	phenotype, the method comprising the steps of:				
3	(a) providing a nucleic acid library comprising nucleotide sequences that				
4	encode at least partially randomized zinc finger proteins;				
5	(b) transducing cells with expression vectors, each comprising a nucleotide				
6	sequence from the library;				
7	(c) culturing the cells so that zinc finger proteins are expressed in the cells,				
8	wherein the zinc finger proteins modulate gene expression in at least some of the cells;				
9	(d) assaying the cells for a selected phenotype and determining whether or				
0	not the cells exhibit the selected phenotype; and				
1	(e) identifying, in cells that exhibit the selected phenotype, the gene or				
2	genes whose expression is modulated by expression of a zinc finger protein, wherein the				
3	gene so identified is associated with the selected phenotype.				
_					
1	2. The method of claim 1, wherein the zinc finger protein has three,				
2	four, or five fingers.				
1	3. The method of claim 1, wherein the library comprises no more than				
2	10 ⁷ clones.				
1	4. The method of claim 1, wherein the cells are physically separated,				
2	individual pools of cells and each individual pool of cells is transduced with an				
3	expression vector comprising a nucleotide sequence from the library.				
1	5. The method of claim 4, wherein the physical separation of the				
2	pools of cells is accomplished by placing each pool of cells in a separate well of a 96,				
3	384, or 1536 well plate.				
	•				
1	6. The method of claim 4, wherein the cells are assayed for the				
2	selected phenotype using liquid handling robots.				
1	7 The method of claim 1, wherein the cells are pooled together and				

1		8.	The method of claim 7, wherein the cells are assayed for the		
2	selected phenotype using flow cytometry.				
1		9.	The method of claim 1, wherein the library is made by finger		
2	grafting, DNA	A shuffli	ing, or codon doping.		
1		10.	The method of claim 1, wherein the zinc finger proteins are fusion		
2	proteins comp	orising a	regulatory domain.		
1		11.	The method of claim 10, wherein the zinc finger proteins are fusion		
2	proteins comp		at least two regulatory domains.		
1		12.	The weeks I of their 10 mb amin the magnitude of amount in palested		
1	ė		The method of claim 10, wherein the regulatory domain is selected		
2		-	sting of a transcriptional repressor, a methyl transferase, a		
3	transcriptional activator, a histone acetyltransferase, and a histone deacetylase.				
1		13.	The method of claim 10, wherein the regulatory domain is VP16 or		
2	KRAB.				
1		14.	The method of claim 1, wherein modulation of gene expression is		
2	repression of				
_	repression or	gene en	p. 400.00		
1		15.	The method of claim 1, wherein modulation of gene expression is		
2	activation of gene expression.				
1		16.	The method of claim 1, wherein the cells are selected from the		
2	group consist	ing of a	nimal cells, plant cells, bacterial cells, protozoal cells, or fungal		
3	cells.	Ü			
1		17.	The method of claim 1, wherein the cells are mammalian cells.		
•		17.	The method of claim 1, wherein the cons are manimum cons.		
1		18.	The method of claim 1, wherein the cells are human cells.		
1		19.	The method of claim 1, wherein expression of the zinc finger		
2	proteins is con	ntrolled	by administration of a small molecule.		
1		20.	The method of claim 19, wherein the small molecule is		
2	tetracycline.				

1

2

1

2

3

1

2

1

2

3

1

2

3

1

2

3

1	21.	The method of claim 1, wherein the expression vectors are a viral			
2	vector.				
1	22.	The method of claim 21, wherein the expression vectors are a			
2	retroviral expression vector, a lentiviral expression vector, an adenoviral expression				
3	vector, or an AAV expression vector.				
1	23.	The method of claim 1, wherein the selected phenotype is related			
2	cancer, nephritis, prostate hypertrophy, hematopoiesis, osteoporosis, obesity,				
3	cardiovascular disease, or diabetes.				

24. The method of claim 1, wherein the zinc finger proteins comprise a Zif268 backbone.

to

- 25. The method of claim 1, wherein genes that are associated with the selected phenotype are identified by comparing differential gene expression patterns in the presence and absence of expression of the zinc finger protein.
- 26. The method of claim 25, wherein differential gene expression patterns are compared using an oligonucleotide array.
- 27. The method of claim 1, wherein genes that are associated with the selected phenotype are identified by using zinc finger proteins from the library of randomized zinc finger proteins to probe YAC or BAC clones.
- 28. The method of claim 1, wherein genes that are associated with the selected phenotype are identified by scanning genomic sequences for target sequences recognized by zinc finger proteins from the library of randomized zinc finger proteins.
- 29. The method of claim 1, wherein genes that are associated with the selected phenotype are identified by cross-linking the zinc finger protein to DNA with which it is associated, followed by immunoprecipitation of the zinc finger protein and sequencing of the DNA.